## WHAT IS CLAIMED IS:

1. A method for analyzing a nucleic acid sample comprised of the steps:

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- (a) forming labeled DNA sample fragments from a nucleic acid sample;
- (b) size separating said sample fragments with size standard fragments, and detecting the fragments to form a sample signal and a size standard signal;
  - (c) transforming the sample signal into size coordinates using the size standard signal; and
  - (d) analyzing the nucleic acid sample in size coordinates.
  - A method for analyzing a nucleic acid sample comprised of the steps:
  - (a) forming labeled DNA sample fragments from a nucleic acid sample;
- 25 (b) size separating and detecting said sample fragments to form a sample signal;
  - (c) forming labeled DNA ladder fragments corresponding to molecular lengths;

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(d) size separating and detecting said ladder fragments to form a ladder signal;

(e) transforming the sample signal into length coordinates using the ladder signal; and

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- (f) analyzing the nucleic acid sample signal in length 5 coordinates.
  - 3. A method as described in Claim 2 wherein after the analyzing step (f) there is the additional step of determining a length or amount of a fragment in the nucleic acid sample.
  - 4. A method as described in Claim 3 wherein after the determining step there is the additional step of finding a gene by positional cloning.
  - 5. A method as described in Claim 3 wherein after the determining step there is the additional step of identifying an individual by DNA profiling.
  - of genetic data comprised of the steps:
  - (a) supplying a software program that automatically scores genetic data;
- 25 (b) forming genetic data that can be scored by the software program;
  - (c) scoring the genetic data using the software program to form a quantity of genetic data; and
  - (d) generating a revenue from computer scoring of genetic data that is related to the quantity.

- 7. A method as described in Claim 6 wherein prior to the step (d) of generating a revenue there are the steps of:
- (e) defining a labor cost of scoring the quantity of genetic data when not using the software program;
  - (f) providing a calculating mechanism for estimating the labor cost from the quantity;
- 10 (g) determining the labor cost based on the quantity; and
  - (h) establishing a price for using the software program that is related to the labor cost.
  - 8. A method as described in Claim 7 wherein the calculating mechanism includes a spreadsheet.

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- 9. A method as described in Claim 7 wherein the calculating mechanism is provided via the Internet.
- 10. A method as described in Claim 7 wherein the calculating mechanism operates interactively via the Internet.
- 25 11. A system for analyzing a nucleic acid sample comprising:
  - (a) means for forming labeled DNA sample fragments from a nucleic acid sample;
  - (b) means for size separating and detecting said sample fragments to form a sample signal, said separating and detecting means in communication with the sample fragments;

- (c) means for forming labeled DNA ladder fragments corresponding to molecular lengths;
- (d) means for size separating and detecting said ladder 5 fragments to form a ladder signal, said separating and detecting means in communication with the ladder fragments;
- (e) means for transforming the sample signal into length coordinates using the ladder signal, said transforming means in10 communication with the signals; and
  - (f) means for analyzing the nucleic acid sample signal in length coordinates, said analyzing means in communication with the transforming means.
  - 12. A method for producing a nucleic acid analysis comprised of the steps:

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- (a) analyzing a first nucleic acid sample on a first size separation instrument to form a first signal;
- (b) analyzing a second nucleic acid sample on a second size separation instrument to form a second signal;
- 25 (c) comparing the first signal with the second signal in a computing device with memory to form a comparison; and
- (d) producing a nucleic acid analysis of the two samples from the comparison that is independent of the size separation 30 instruments used.
  - 13. A method as described in Claim 12 wherein the size separation instrument is a DNA sequencer that uses electrophoresis.

- 14. A method as described in Claim 12 wherein the nucleic acid analysis characterizes a size or amount of DNA in one of the nucleic acid samples.
- 15. A method as described in Claim 12 wherein the nucleic acid analysis finds a gene by positional cloning.

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- 16. A method as described in Claim 12 wherein the nucleic acid analysis identifies an individual by DNA profiling.
  - 1/2. A method for resolving DNA mixtures comprised of the steps:
    - (a) obtaining DNA profile data that include a mixed
    - (b) representing the data in a linear equation;
    - (c) deriving a solution from the linear equation; and
    - (d) resolving the DNA mixture from the solution.
- 18. A method as described in Claim 17 wherein the
  25 obtaining step (a) includes the step of performing a PCR on an STR locus of an individual.
- 19. A method as described in Claim 17 wherein the representing step (b) includes a matrix or vector representation.
  - 20. A method as described in Claim 17 wherein the deriving step (c) includes an optimization procedure.

21. A method as described in Claim 17 wherein the deriving step (c) includes a matrix operation.

Signature  $\lambda$  3 | 22. A method as described in Claim 17 wherein the 5 resolving step (d) produces the genotype of an individual.

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